IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Application of

Applicant(s) : Catherine M. Verfaillie et al.

Application No. : 10/561,826

Filed : October 17, 2006

Title : Neuronal Differentiation of Stem Cells

Examiner : Chang Yu Wang

Art Unit : 1649 Attorney Docket : 890003-2006.1

Attorney Docker : 050005-2000.1

DECLARATION UNDER UNDER 37 C.F.R. § 1.132

Commissioner for Patents P. O. Box 1450 Alexandria, VA 22313-1450

Sir:

The undersigned, Catherine M. Verfaillie, Ph.D., declares and states:

I am Professor of Medicine and Director Stem cell Institute, KULeuven, Leuven I am a co-inventor on the above-captioned patent application.

I am the subject of the attached *Curriculum Vitae* and author of the publications listed on the attachment to the *Curriculum Vitae*. On the information and facts contained in those documents, I submit that I am an expert in the field of Stem cell research. In view of these credentials, I believe that I am qualified to speak on the skill and knowledge of the person of ordinary skill in these fields.

I have read and understand the subject matter of the above-captioned patent application. I have read the first Office Action, dated March 17, 2008, and the second Office Action, dated December 24, 2008. I have read and understand the references cited by the Examiner to support the rejections in the Office Actions. These include U.S. 2003/0211605 to Lee et al. ("Lee") and WO 02/086073 to Studer et al. ("Studer").

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It is my opinion, based on the scientific evidence and reasoning below, that the rejections are based on an

incorrect assumption about (1) how cells should respond to basic fibroblast growth factor (bFGF),

fibroblast growth factor 8 (FGF8), Sonic Hedgehog (SHH), and brain-derived neurotropic factor (BDNF)

when these are administered simultaneously as opposed to when these are administered sequentially, and

(2) the cell type being acted upon by bFGF, FGF8, SHH, and BDNF in the cited art as opposed to the

(a) the contribution of area, area, and area area area area.

claimed method.

In the earlier Office Action, on page 6, the Examiner rejects the claims as being obvious over Studer in

view of Lee. In the later Office Action, the Examiner provides particular reasons for this rejection.

Specifically, the Examiner appears to believe that one would generally expect the same result whether one

adds all factors simultaneously or sequentially. Specifically, the Examiner states on page 7 of the Office

Action, "... at the end of the final steps, the culture medium contains the identical growth factors as those

in Studer's to induce neuronal differentiation." The Examiner then reasons, "... because at the end the

culture medium still contains the same growth factors and the same cultured ES cells, ...would be

induced to differentiate into neurons." (Emphasis added.) Essentially, the Examiner appears to take the

position that sequential exposure should have no significant effect because the factors and the cells

exposed to them are the same. I do not agree with this reasoning for the reasons that follow.

The End Product of Sequential Exposure to Factors Cannot be Reasonably Predicted Based on the Results

of Simultaneous Exposure to the Same Factors

When one introduces a cocktail of factors, as Studer and Lee have done, all the factors are exposed at

once to one discrete cell type as to functional, transcriptional, translational, and morphological

characteristics. Thus, the factors are acting at the <u>same time</u> on the <u>same cell</u>. Such exposure produces a

certain end product. But, in the claimed methods, there are three phenotypically discrete cell types that

are being acted upon: (1) the starting cells, (2) the cells that have been exposed to bFGF but not to FGF8

and SHH, and (3) the cells that have been exposed to bFGF, FGF8 and SHH, but not BDNF. Each

mitogen would have a specific effect on the cell to produce a phenotypically discrete cell type. So, if one

exposes a cell to bFGF, FGF8, SHH, and BDNF at the same time, the FGF8/SHH is not acting on a cell with a phenotype created by exposure only to bFGF; and the BDNF is not acting on a cell with a phenotype created by exposure only to bFGF, SHH, and FGF8. Therefore, contrary to what the Examiner asserts, when all the factors are in the medium, they are <u>not</u> acting on the "same cultured ES cells." Because of this, one cannot reasonably predict that the result will be the same as the result obtained by Studer and/or Lee.

Although we have not done an experimental comparison between the Studer and Lee end products and the end products obtained using our own steps a) through c), I will discuss an application of the principle illustrated with a differentiation protocol that we conducted in our laboratory pertaining to differentiation of adult bone marrow stem cells into functional hepatocyte-like cells (Snykers et al., Toxicological Sciences, 94:330-341 (2006)). Although the reference is directed to differentiation of stem cells into hepatocytes, the principle applies: sequential exposure to factors can result in quite a different end product than simultaneous exposure ("cocktail").

As reported in this reference, we compared the end product obtained by using a cocktail of factors versus sequential exposure to the factors. I will not go into great detail about this reference as the Examiner is fully qualified to assess the reference. However, I will briefly give an outline of the rationale and the results. It had been previously been shown that bone marrow stem cells could differentiate into hepatocyte-like cells from a simultaneous exposure to a mixture of cytokines and growth factors. To try to improve the end product, the cells were exposed to the same factors in a sequential way. Characterization of the cells over a period of time and after exposure to each factor were characterized in several ways: (1) morphology; (2) mRNA expression of hepatocyte-specific genes; (3) protein expression of hepatocyte-specific genes; and (4) hepatic functionality as assessed by albumin secretion, ureogenesis, glycogen storage, and CYP protein expression activity and inducibility.

Our results showed the following. With respect to morphology, using the sequential procedure, the stem cells acquired morphological features similar to those of primary hepatocytes, particularly polygonalshaped and bi-nucleated cells. In contrast, using the previous approach, a heterogenous population of

epithelioid cells and other cell types was obtained with no polygonal-shaped cells and only a few bi-

nucleated cells. With respect to liver associated genes and proteins, more than 85% of these epithelioid

cells expressed these genes and expressed them in a comparable time-dependent manner as observed

during in vivo liver embryogenesis. In contrast, with the cells exposed to the cocktail, the expression

patterns differed from the normal sequence in that HNF1α expression preceded that of albumin. In

addition, significantly lower levels of liver-specific markers were expressed. With respect to functional

maturation, this occurred with both experimental protocols but to a different extent. Hepatic metabolic

functions, including albumin secretion, urea production, etc., were manifested most prominently upon

sequential exposure to hepatogenic factors.

This illustrates the principle that the end product of sequential exposure to factors cannot be reasonably

predicted based on the results of simultaneous exposure to the same factors.

My understanding is that the rejection of obviousness must be based on motivation to change the

Studer/Lee procedure. It is my opinion that one would not have been motivated to alter the procedure of

Studer and/or Lee because they would not have reasonably expected to produce the same result. Further,

my understanding is that the rejection of obviousness must include a reasonable expectation that the same

end product would be obtained. As I understand it, another way of looking at this is that it must be

reasonably predictable that the same end product would be produced by simultaneous and sequential

exposure to the factors. The Examiner seems to assert that position. But, it is my opinion as an expert in

the field, that it was not reasonably predictable that the end product in the prior art would be produced by

sequential exposure to the factors.

Summary

In the Studer and Lee method, one discrete cell type (neuronal commitment) is exposed to all three factors

at once to create a discrete, functional, morphological, and transcriptional and translational profile. Using

the sequential method, however, the starting cells are first exposed to bFGF, which creates a cell with a

specific morphology, function, and transcriptional and translational profile. It is this cell that is acted

upon by SHH and FGF8, not the original starting cell as in the "cocktail" method. Then, exposure to

FGF8 and SHH produces a cell with a second discrete type of morphology, function, and transcription

and translational profile. It is this cell that is acted on by the BDNF, not the original starting cell as in the

"cocktail" method. That is what makes the end product not reasonably predictable.

Studer/Lee Do Not Apply the Factors to Multipotent Cells

I also point out that Studer/Lee do not apply bFGF to embryonic stem cells, i.e., to a multipotent stem

cell. Studer/Lee apply bFGF to a cell already committed to a neural fate. Studer/Lee form embryoid

bodies from embryonic stem cells and grow these embryoid bodies without any differentiation factors,

selecting for the cells that have undergone neural commitment. It is those cells that have undergone

neural commitment that are exposed to the mitogen(s), SHH, and FGF8. The person of ordinary skill

would have expected that neural commitment was needed prior to the application of mitogen(s) SHH and

FGF8.

In contrast, in the claimed methods, bFGF is applied directly to multipotent stem cells and not to a

neurally committed cell. In fact, it is the bFGF that induces neural commitment in these cells.

For this reason alone, I believe that the claimed method would not motivate one to practice the claimed

method.

CONCLUSION

It is my opinion, based on the scientific evidence and reasoning set forth above, that the rejection lacks

sufficient scientific basis for finding obviousness.

In my opinion, the person of ordinary skill in this field would not have been motivated to drastically

change the approach of Studer and/or Lee from a cocktail approach to a sequential approach and would

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not have had a reasonable expectation that the same results would be obtained if they did change the

approach.

The Studer/Lee method applied to a neural committed cell, not an embryonic stem cell; therefore, the

person of ordinary skill could not have reasonably expected successful application of the Studer/Lee

method (i.e., exposure to mitogen(s) FGF8 and SHH) unless embryonic stem cells were committed to a

neural fate before the factors were applied.

All statements made herein of my own knowledge are true and all statements made on information

believed to be true. These statements were made with the knowledge that willful false statements and the

like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the

United States Code and that such willful false statements may jeopardize the validity of the application or

any patent issued thereon.

6/1109

Date

Catherine M. Verfaillie, Ph.

Curriculum vitae: Catherine Maria Verfaillie Januari, 2009

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CITIZENSHIP: Belgium

Permanent Resident USA

EDUCATION: M.D.: U. of Leuven Medical School, Leuven, Belgium, 1975-1982

POST-GRADUATE TRAINING:

Internship: 1982-83, AZ St Jan Hospital, Brugge, Belgium

Residency: 1983-85, U. of Leuven, Belgium

Fellow in Hematology: 1985-87, U. of Leuven, Belgium

Post-doctoral Fellow: 1987-89, U. of Minnesota, Minneapolis, MN

PROFESSIONAL APPOINTMENTS:

Instructor, Dep. of Medicine, University of Minnesota, 1989-90,

Assistant Professor of Medicine, University of Minnesota 1991-1995

Associate Professor of Medicine, University of Minnesota 1995-1998

Professor of Medicine, University of Minnesota 1998-

Director Stem Cell Biology Program, University of Minnesota, 1996-1999

Director, Stem Cell Institute, University of Minnesota, 1999-2006

Buitengewoon Hoogleraar, Katholieke Universiteit, Leuven, 2005-

Director, Stamcel Instituut, Katholieke Universiteit, Leuven, 2005-

Member, BME Graduate Program, since 1992

Member, MICaB Graduate Program, since 1992 Member, MD/Ph.D Graduate Program, since 1994

Member Cancer Center, 1994

Member, GCD Graduate Program, since 2000

Member, Clinical Laboratory Science Graduate Program, since 2000

Member, Neuroscience Graduate Program, since 2002

Member Doctoral School, Department of molecular and Cellular Therapy, K.U.Leuven

CERTIFICATION:

FMGEMS-1987 FLEX-1990

HONORS:

M.D., Summa Cum Laude, 1982

Special Fellow, Leukemia Society of America, 1991

Special Fellow, 'Fundacion Internacional Jose Carreras Para La Lucha Contra La Leucemia ',1991

Young Investigator Award, International Society of Exp. Hematology, 1992

Scholar, Leukemia Society of America, 1995

Outstanding Investigator Award, Central Society, 1996

Elected, Member, American Society of Clinical Investigation, 1996

Tulloch Chair in Stem Cell Biology, Genetics and Genomics, 1999

Anderson Chair in Stem Cell Biology, 1999

Elected, Councilor, American Society of Clinical Investigation, 2001

McKnight's Presidential Chair in Stem Cell Biology, 2001

Elected, Member, American Association of Professors, 2003

Vice President, International Society of Experimental Hematology, 2002

President Elect, International Society of Experimental Hematology, 2002 President, International Society of Experimental Hematology, 2004

4th Annual Landazuri Award, University of Navarra, Pamplona, Spain 2002

Damashek Medal, American Society of Hematology, 2002

Honorary Doctorate, Katholieke Universiteit, Belgium, 2003

Distinguished Woman Scholar Award, University Of Minnesota, 2003

2003 Jose Carreras Award, European Society of Hematology, 2003 Forum Engelberg Prize, Lucerne, Switzerland, 2003

Gulden Spoor voor Vlaamse Internationale Uitstraling, Vlaanderen-Europa, 2003

Star Award, Minnesota Hematology Oncology, 2004

Stewart-Niewiarowski Award for Women in Vascular Biology. 2004

Jimenez Diaz Price for Scientific Achievements in Research, 2004

Vlerick Award, 2005

Honorary Member, BeWiSe, 2005

Gabriella Moortgat Prijs, 2006

Bijzonder Hoogleraar, TEFAF Oncology Wisselleerstoel, 2009, Faculty of Health, Medicine, and Life Sciences, Universiteit van Maastricht

Cariplo Professor, University of Pavia, Italy, 2009-2011

NAMED LECTURES:

Presidential Symposium, American Society of Hematology, San Francisco, 2000 Nobel-Forum Lecture series, Karolinska Institute, Stockholm, Sweden, 2000 Linda Laubenstein Memorial Lecture, NYU, New York, 2000 Clement Finch Visiting Professor, University of Washington, Seattle, 2002

4th Annual Landazuri Lecture, University of Navarra, Pampiona, Spain, 2002

Gerhard Smith Memorial Lectureship, City of Hope, CA, 2002

Fiftieth Chalmers J Lyons Memorial Lectureship, AAOMS, Chicago, IL, 2002

Forum Engelberg Award Seminar, Lucerne, Switzerland, 2003

Hohenberg Lecture, university of Pennsylvania, PA, 2003

Kilo Professorship, Washington University, St. Louis, MO, 2003

Evans Lecture, Boston University, Boston, MA, 2003 Stewart-Niewiarowski lecture, Temple University, Boston, MA, 2004

KeKuLe Lecture 2004, Antwerp, Belgium, 2004

NIH Director's Lecture, Washington, DC, 2004

Bendit Lecture, University of Washington, Seattle, WA, 2004

Jimenez Diaz Commemorative Lecture, Madrid, Spain, 2004

Presidential Symposium, ASGT, Minneapolis, MN, 2004

Latta Lecture, University of Nebraska, Omaha, 2005

Presidential Symposium, Am Soc for Investigative Pathology, San Diego, 2005

Moloney Lecture, Brigham & Women's Hospital, Boston MA, 2005

Ada Comstock Inaugural Lecture, Minneapolis, MN, 2005

Brecher Lecture, San Francisco, CA, 2005

Visiting Professor Department of Medicine, Vanderbilt University, 2006

Presidential Symposium, ESTRO, Leipzig, Germany, 2006

Pierre Stryckmans Memorial Lecture, Brussels 2007

PROFESSIONAL ASSOCIATIONS:

American Federation for Clinical Research (AFCR) American Society of Gene Therapy (ASGT)

American Society of Hematology (ASH)

American Society of Stem Cell and Bone Marrow Transplantation (ASBMT)

International Society for Hematotherapy and Graft Engineering (ISHAGE)

International Society of Experimental Hematology (ISEH)

International Bone Marrow Transplantation Registry (IBMTR)

Autologous Bone Marrow Transplantation Registry (ABMTR) International Society for Stem Cell Research (ISSCR)

COMMITTEE ASSIGNMENTS (Extramural):

Treasurer, International Society of Experimental Hematology, 1997-2001 Councilor, American Society of Clinical Investigation, 2001-2005

Board of Directors, American Society of Blood and Bone Marrow Transplantation, 2001-2005

Councilor, Society Cell transplantation, 2002-2006

Member, Scientific Committee of the European School of Hematology, 2004-

Chair, Policy Committee, ISSCR, 2002-2006

Member, Scientific Subcommittee on Transfusion Medicine, ASH, 1996-1998

Member, Scientific Subcommittee on Growth Factors, ASH, 1998-2002

Member, Awards Committee, ASH, 2003-2007

Member, Publications Committee, ISEH, 2001-2005

Member, Committee on Hemopoletic Cell and Gene Therapy, ASGT, 98-02

Member, Stem Cell Evaluation Committee, ISHAGE, 1997-2001

Member, Mesenchymal Stem Cell Committee, ISHAGE, 1999-2004

Member, Stem cell Expansion Committee, ISHAGE, 2001-2003

Chair, VA Merit Award Heme Study Section, 2001-2002

Member, NIH study section, Heme I, 2000-2004

Member, VA Merit Award Heme Study Section, 1999-2002

Member, LSA Translational Awards Review Committee, 1999-2004

Member, Telethon Scientific Committee, Italy, 2001-2005 CIRM review committee, 2006-

Review committee, 2006-

Member CNRS review panel, Belgium, 2008-2012

Ad hoc reviewer, Juvenile Diabetes Research Fund

Ad hoc reviewer, Muscular Dystrophy Association

Ad hoc reviewer, Leukemia Research Fund, Great Britain

Ad hoc reviewer, Welcome Trust, Great Britain

Ad hoc reviewer, Research Council, Canada

Ad hoc reviewer, Nationaal Fonds Wetenschapelijk Onderzoek, Belgium

Ad hoc reviewer, Associazone Italiana per la Ricerca Sul Cancro

Ad hoc reviewer, Israel Science Foundation Ad hoc reviewer, Dutch Cancer Society

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Ad Hoc reviewer, Medical and Health Services Research Division, Ireland.

Ad hoc reviewer, European Commission, FP6

Ad hoc reviewer, European Commission, FP7

Ad hoc reviewer, European Research Council, 2007-

COMMITTEE ASSIGNMENTS (Intramural):

Promotion and Tenure Committee, Department of Medicine 1997-2001

Research Committee, Department of Medicine, since 1997

Steering Committee MD/Ph.D. Program, 1998-2005

Molecular Medicine Planning Committee, 1998-2000
AHC, Functional Genomics Advisory Group, 2000-2004

Consortium on Law and Values in Health, Environment & the Life Sciences, 2001-6

CONSULTANT/ADVISOR

Member International Scientific Advisory Board, UK Government and Wellcome Trust Joint Infrastructure Fund,

Member Advisory Committee, Institute of Hematology, Chinese Academy of Sciences & Peking Union Medical College, 2000-2005

Member, Advisory Committee Tissues of Life Project, Science Museum of Minnesota, 2000-2006

Member Scientific Advisory Panel, University of Nebraska Stem Cell Biology Research Center, 1999-2004

Member Scientific Advisory Panel for National Stem Cell Resource, Corriel Institute, Camden, NJ, 2000-2004

Member, Advisory Committee, Mayo Clinic Myeloma PO1, 1998-2004

Member, Stem Cell Advisory Committee, National Research Institute, Taiwan, 2002-

Consultant, Athersys Inc., Cleveland, OH, 2002-

Member, Advisory Board, Center for Transgene Therapy and Gene Therapy, VIB, Leuven, Belgium, 2003
Member Advisory Panel. Stem Cell GAP, NIH, 2003-5

Member Advisory Panel, Stem Cell GAP, Nin, 2005-

Member, Scientific Advisory Board, Oncostem Therapeutics, Salamanca, Spain, 2004-2007

Member, Scientific Advisory Board, Toronto McLaughlin Centre, 2004-2007

Member, Scientific Advisory Board, Framework 6 Program Beta Cells, 2004

Member, Scientific Advisory Board, DPTE, 2005-

Member, Scientific Advisory Board, Case Western University Stem Cell Institute, 2005-2008

Lid Raad van Advles, EOS, 2005-

Member Advisory Committee, Itinera, 2006-

Advisory Board Regenerative Medicine, the Netherlands, 2007-2009

Co-Chair Research and Quality Assurance Evaluation, Lund University, 2008

Member, Review Committee Science Foundation Ireland REMEDI CSET, 2008

Member, Scientific Advisory Committee, Fondazione Roma, 2008-Member, Advisory Board, FP7 Infarct Cell Therapy project, PLE Hofer, 2008-

Member, Advisory Board, FP7 Intarct Cell Therapy project, PLE Hoter, 2008-Member International Advisory Board Norway Stem Cell Center, 2008-

Member Advisory Board, EC project "Infarct Cell Therapy", 2009-

ASSOCIATE EDITOR:

Experimental Hematology, 2003-2008

Experimental Hematology, 1998-200

Leukemia, 1997-2002

Hematologia, Citocinas, Immunoterapia Y Terapia Cellular, 1997-2002

Stem Cell Reviews, 2004-2008

PloS-1, 2008-

EDITORIAL BOARD:

Blood, 1995-1999

Experimental Hematology, 1996-1998

Leukemia, 1996-1997

Cytotherapy, 1999-2003

Journal of Biology of Blood and Marrow Transplantation, 2001-2005

Cloning & Stem Cells, 2001-2005

Current Gene Therapy, 2005-

Journal of Engineering and Regenerative Medicine, 2006-

Stem Cells, 2007-2008

Stem Cells International, 2008-

PATENTS:

WO9718298: Ex vivo culture of stem cells

CA2381292: Multipotent adult stem cells and methods for isolation

US2007022482: High-throughput functional analysis of gene expression

WO2006086639: Vascular/lymphatic endothelial cells

WO2006047743: Swine multipotent adult progenitor cells

WO2005045012: Endodermal stem cells in liver and methods for isolation thereof

WO2005003320: Neuronal differentiation of stem cells

WO2008063675: Endodermal progenitor cells

US2002081733: Method to prepare drug-resistant, non-malignant hematopoietic cells

WO2004050859: Homologous recombination in multipotent adult progenitor cells

WO9513088A1: Stroma-derived stem cell growth factors

WO9320184: Method for culturing hematopoietic cells

WO2002040718: Method to identify genes associated with chronic myelogenous leukemia

AU2006304318; Differentiation of non-embryonic stem cells to cells having a pancreatic phenotype

AU2005331534: Use of MAPC or progeny therefrom to populate lymphohematopoietic tissues

US11/808933: High Oct3/4 MAPCs and methods therefor

US 61/022121: Stem cell aggregates and methods for making and using

US08/82108: Optimized methods for differentiation of cells into cells with hepatocyte and hepatocyte

progenitor phenotypes, cells produced by the methods, and methods for using the cells

US - GB 0822483.4: Maintenance/expansion of HSCs

US 60/690089 *: HSC Self-Renewal (CIP of US2007022482)

TRAINEES:

High School Students		Current Position	
Evan Cobbs	2002-2003	U of Madison, undergraduate	
Nicole Ali	2002-2004	Harvard University, undergraduate	
Ricky Jones	2002-2004	CalTech, undergraduate	
Sam Bjork	2003-2005	Harvard University, undergraduate	

Undergraduate Students

Venita Chandra 1997-1998 Medical School, U of Chicago

Current Position

Sarah Aldrich	2000, 2001	Medical School, U of Chicago
Aaron Lisberg	2001, 2002	MD/PhD program,
Zach Kastenberg	2002-2004	Medical School, U. of Minnesota
Eric Rarhman	2002-2005	Graduate School, U of Minnesota
Lee Sandquist	2003-2005	Medical School, U of Minnesota
Juliana Hagenbrock	2003-2006	PhD Student, University of Amsterdam, The Netherlands
April Breyer	2003-2006	Law School, Boston College Law School
Jennifer Gravelle	2003-2005	Medical School, U of Minnesota
Mike Felten	2003-2005	Medical School, U of Minnesota
Thomas Szynski	2003-2005	Medical School, Harvard U
Medical Student Advisor		Current Position

Sjoban Keel, B.S. Troy Lund, Ph.D.	1997-1998	Fellow, Hematology, U. of Washington
David Dyle, B.S.	1998-2002	Pediatric Heme-Onc Fellow, U of Minnesota
	1998-2000 2003-2005	Fellow, Genetics, U. of Washington
Zubaid Rafique, B.S.		
Tzu-Fei Wang. M.S. Eleanor Chen, Ph.D	2004-2005 2004-2006	Pathology Rosidancy Hannard II
Eleanor Chen, FILD	2004-2006	Pathology Residency, Harvard U
Medical Resident Research		Current Position
Jade Anderson, M.D.	1999-2000	Fellow, Hematology, U. of Minnesota
Ken Lee, M.D.	2000-2001	Fellow, Cardiology, UCSF
Modiani Davidana Adulasa		Command Barathian
Medical Resident Advisor		Current Position
Sjoban Keel, M.D.	2000-2001	Fellow, Hematology, U. of Washington
David Dyle, M.D.	2000-2004	Fellow, Genetics, U. of Washington
Masters Students		Current Position
Yuehua Jiang, M.D.	1997-1999	Assist. Prof., U of Minnesota
Sofia Melikova, B.S.	2000-1002	Scientist, R&D Systems, Minneapolis
Ben Vaessen, B.S.	2000-2002	Scientist, Glaxo-Smith-Kline
Avinash Jayaswal, B.S.	2002-2003	Medical School, UCL, Brussels, Belgium
Qing Cai	2006-2007	PhD Student, K.U.Leuven
Janick Beckers	2006-2007	PhD Student, K.U.Leuven
David Zwaenepoel	2006-2007	Scientist, ReGenesys, BVBA
Adriaan Campo	2006-2007	PhD Student, U. Antwerpen
Maria Aelberts	2007	PhD Student, U. of Copenhagen
Antonio LoNigro	2007-2008	PhD Student, K.U.Leuven
Simone Calzolari	2007-2008	PhD Student, U. of Barcelona
Kim Van Uytsel	2007-2008	PhD Student, K.U.Leuven
Jasper Wouters	2007-2008	PhD Student, K.U.Leuven
Tine Verryckt	2007-2008	
Olivier Govaere	2007-2008	PhD Student, K.U.Leuven
Lotte Vanbrabant	2007-2008	Technician, K.U.Leuven
Vijay Kumar	2008	PhD Student, U of Toronto
Alessandra Familiari	2008-2009	
Caterina DiPrieto	2008-2009	
Graduate Students:		Current Position
Beverly Lundell, MS.	1993-1996	Scientist, Aastrom, Michigan
Eugene Liu, M.D.	1996-2000	Assist Prof of Medicine, U. of Taipei, Taiwan
Scott Dylla, M.S.	1998-2002	Post-doc, Irving Weissman, Stanford U
Fernando Ulloa, B.S	2002-2006	Scientist, Glaxo Smith Kline
Lucas Chase, B.S.	2003-2006	Scientist, Invitrogen
Eric Mendenhall, B.S.	2003-2006	Post-doc, Bernstein lab, MIT
Lepeng Zeng, B.S.	2003-2006	Scientist, Medtronic
Jeff Ross, B.S., M.S.	2003-2006	Scientist, Surmodics
Ben Kidder, B.\$.	2003-2007	Postdoc, Serono, Boston
Shannon Buckley, B.S.	2004-2009	
Annelies Crabbe, BS	2006-	
Valerie Roobroeck, BS	2006-	

Kartik Subramanian, BS	2006-	
Yonsil Park, BS	2006-	
Jason Owens, BS	2007-	
Qing Cai, BS, MS	2007-	
Kim Van Uytsel, BS, MS	2008-	
Antonio LoNigro, BS, MS	2008-	
Rojin Abraham MD	2008-	
Elda Mineola, MD	2008-	
, ···-		
MD/PhD students:		Current Position
Morayma Reyes, B.S.	1997-2001	Ass. Prof. Lab Medicine, U of Washington, Seattle
Robert Schwartz, B.S.	2000-2004	Fellow Gastroenterology, Harvard University, Boston
Craig Eckfeldt, B.S.	2002-2005	Resident Internal Med, U. of Minnesota
Sarah Frommer, B.S.	2002-2006	Medical School, U of Minnesota
Terri Burns, B.S.	2003-2007	Medical School, U. of Minnesota
Postdoctoral Fellows		
Chunjin Ding, M.D.	1992-1994	Scientist, Ely Lilly, Indianapolis
Huilin Qi, Ph.D.	1998-2003	Scientist, Yale University
Dean Aguiar, Ph.D.	1999-2001	Staff Scientist, Pharmacia
Yuehua Jiang, MS, MD	1999-2002	Assist. Prof., U of Minnesota
Stephanie Salesse, Ph.D.	2000-2005	Assist. Prof, U of Reims, France
Mo Dao, Ph.D.	2001-2004	Post Doc, N Taylor, Montpellier, France
Troy Lund, MD, Ph.D.	2002-2003	Assist. Prof, Department of Pediatrics, U of Minnesota
Yves Heremans, Ph.D.	2002-2005	Instructor VUB, Belgium
Uma Lakshmipathy, Ph.D.	2002-2005	Staff Scientist, Invitrogen, CA
Beatrice Pelacho, Ph.D.	2002-2005	Instructor, U of Navarra, Pamplona, Spain
Aernout Luttun, Ph.D.	2002-2006	Assist. Prof, K.U.Leuven, Belgium
Miguel Barajas, Ph.D.	2004-2006	Assist. Prof., U of Navarra, Pamplona, Spain
Marta Serafini, Ph.D.	2004-2006	Assist. Prof., Fondazione M.Tettamanti M.De Marchi
		Onlus, Monza, Italy
Rik Snoeckx, Ph.D.	2005-2009	Postdoctoral fellow, J Cools lab, VIB-KULeuven
Kris Van den Boogaert, Ph.D.	2005-	
Carlos Clavel, Ph.D.	2005-2008	Postdoc, Einsteln University, USA
Fernando Ulloa, Ph.D.	2006-2008	Scientist, Glaxo Smith Kline
Martine Geeraerts, Ph.D.	2006-	
Pau Sancho-Bru, Ph.D.	2007-2009	
Takeshi Shimizu, Ph.D.	2007-	
Jeroen DeClercq, Ph.D.	2007-	
Bipasha Bose, Ph.D.	2008-2009	
Anujit Kumar, Ph.D.	2008-	
Satish Kumar, Ph.D.	2009-	
Yong Li, Ph.D.	2008-	
Fellows		Current Position
Jeffrey S Miller, M.D.	1991-1994	Prof. of Medicine, Director, Translational Research
, •,		Program, Cancer center, U. of MN
Pankaj Gupta, M.D.	1992-1995	Prof. of Medicine, U. of MN
Randolph Hurley, M.D.	1992-1995	Staff Physician, Health Partners, St Paul, MN
Ravi Bhatia, M.D.	1992-1996	Prof. of Medicine, Director Stem Cell Program, City of Hope
K.Y. Chiang, M.D., Ph.D.	1993-1995	Assist Prof. Pediatrics, Emory University, Atlanta GA
Vivek Roy, M.D.	1994-1996	Assist. Prof. of Medicine, Mayo Clinic, FL
Robert Zhao, M.D., Ph.D	1995-1998	Professor, Professor, Chinese Acad. of Sci. & Peking
,		Union Medical College
Michael Punzel, M.D.	1996-1998	Assoc Prof., U. of Duesseldorf, Germany
Felipe Prosper, M.D.	1995-1997	Prof. of Medicine and Director Stem Cell Program, U. of
		Navarra, Pamplona, Spain
Michel Delforge, M.D.	1995-1996	Assoc. Prof, KULeuven, Belgium
Juliet Barker, M.D.	1997-1999	Assoc. Prof. of Medicine, Sloan Kettering Institute, NYC
Ian Lewis, M.D., Ph.D.	1997-1999	Assoc. Prof. of Medicine, U. of Adelaide, Australia
Chris Lamming, M.D.	1999-2002	Pediatric Fellow, Baylor College, Houston, TX
Claudio Brunstein, M.D.	1999-2002	Assist Prof. of Medicine, U of Minnesota

Koen Theunissen, M.D. 1999-2001 Hematologist, Virga Jesse Hospital, Hasselt Assist Prof. of Medicine, U of Minnesota Balkrishna Jahagirdar, M.D. 2000-2002 Eugene Liu, M.D., Ph.D. 2000-2002 Assist Prof. of Medicine, U. of Taipei, Taiwan 2001-2003 Instructor Hematology, U. of Rotterdam, NL Moica Jongen, M.D. 2001-2003 Assist. Prof, U of Honk-Kong Anskar Leung, M.D., Ph.D. Masayuki Oki, M.D. 2003-2006 Assist, Prof. Tokai U. Tokio, Japan 2004-2007 Consultant Haematologist, St James's Hospital and Catherine Flynn, M.D. Coombe Women's Hospital, Dublin Karen Pauwelyn, M.D. 2005-2009 Fellow, Hepatology, K.U.Leuven Helene Schoemans, M.D. 2006-2008 Fellow, Hematology, K.U.Leuven 2007-

INVITED LECTURES (Since 2007)

Philip Roelandt, M.D.

2007

Keynote Speaker, Basel Switzerland

First Connecticut International Stem Cell Symposium, Hartford, CT

Grand Challenge Meeting 4, Lugano, Switzerland

Symposium on tissue reconstruction, UCL, Brussels, Belgium

8th Advanced Summer Course in Cell - Materials Interactions, Instituto de Engenharia Biomédica, Porto, Portugal

MSC2007, Adult Mesenchymal Stem Cells in Regenerative Medicine, Cleveland, OH

The Second UK Mesenchymal Stem Cell Meeting, U of York, Great Britain

International Symposium "Stem cells, Development and Regulation" Amsterdam, The Netherlands

Symposium on Cardiovascular Regenerative Medicine, NIH, Washington DC, USA

Spanish National Congress of Surgery, San Sebastian, Spain

Scottish Stem Cell Network, Glasgow, Scotland

Interhospital Endocrine Rounds, Montreal, Canada

Research Seminar, Buck Institute, CA

Research Seminar, U Liege, Belgium

Research Seminar, VUB, Brussel

2008

Keynote Lecture, EPISTEM conference, Gent, Belgium

Kevnote Lecture, VIB, Blankenberge, Belgium

Keynote Lecture, Italian Society for Biotechnology and Medical Engineering, Rieti, Italy

Keynote Lecture, Wadden Symposium on Diabetes, Texel, the Netherlands

Keynote Lecture, Annual Norwegian Stem Cell Network, Oslo Norway

Keynote Lecture, Itera Conference, Maastricht, the Netherlands

Keynote Lecture, Annual science day of the GROW, Maastricht, The Netherlands EuroSTELLS, Stem Cell Niche Meeting, Barcelona, Spain

Symposium "Pluripotency and differentiation in embryos and stem cells", Pavia, Italy

The Adult Stem/Progenitor Cell Niche, Brussels, Belgium

Nederlandse Vereniging voor Hematologie, Papendal, the Netherlands

Annual Wound Healing Society Meeting, San Diego, USA

International Stem Cell Meeting, Tel Aviv, Israel

EAE/ESH 2nd symposium on MSC, Mandelieu, France

Summer School, Barsinghause, Germany

Nobel Forum Cancer Stem Cell Conference, Stockholm, Sweden

4th International Conference on Regenerative Hepatology, Dusseldorf, Germany

European Society of Gene Therapy, Brugge, Belgium

Mayo Clinic Stem Cell and Regeneration Symposium, Rochester, MN, USA

The 5th Dubai International Conference for Medical Sciences, Dubai

EU-EPC Roundtable, Brussels Belgium

2009

Plenary Lecture, Annual Meeting, French Society of Blood Transfusion, Strasbourg, France

Plenary Lecture, Annual Meeting, ESGT, Hannover, Germany

The Sanguin Spring Seminar, Amsterdam, the Netherlands

Flanders Bio Seminar Tissue Engineering, Brussels

Research Seminar, UCL, Brussels, Belgium

Research Seminar, U Frankfurt, Frankfurt, Germany

Epiplastcarcinoma Marle Curie RTN network meeting, Leuven, Belgium

MEETING ORGANIZER

Yearly Meeting ISEH, 2005, Glasgow, Scotland

Yearly meeting ISEH, 2006, Minneapolis, MN

Keystone meeting, Stem Cells, 2006, Whistler, Canada

Mesenchymal Stem Cells (1), 2006, Mandelieu, France

Mesenchymal Stem Cells (2), 2008, Mandelieu, France

Muscular Dystrophy Symposium Leuven, October 2, 2008

Stem Cells: Biology and Applications; Sponsored by FP6-STROKEMAP and TEFAF, Leuven, 2009

RESEARCH INTERESTS:

1. NORMAL HEMATOPOIESIS:

- Regulation of normal human hematopoietic stem cell proliferation, differentiation and lineage commitment by
- cytokines and components of the extracellular matrix using in vitro as well as in vivo xenogeneic transplant models.

Molecular characterization of hematopoietic stem cells by functional genomics, and zebrafish model of hematopolesis PLURIPOTENT STEM CELLS (MAPC, ESC, IPS).

- Purification, expansion and characterization of differentiation to mesodermal, ectodermal and endodermal lineages
- Characterization of molecular determinants of pluripotent and multipotent stem cell phenotype, and of dedifferentiation and differentiation
- Evaluation of therapeutic potential in congenital disorders or for the treatment of vascular, neurodegenerative disorders, hepatic disorders and diabetes.

CURRENT FUNDING:

- PO1-CA-65493-06 (PJ P McGlave): Biology and Transplantation of Human Stem cells Project Period: 7/1/2000 -6/30/2005, Project Leader: project 1; Annual Direct Cost: \$210,000/year
- FWO (PI Verfaillie) Het potentieel van multipotente adulte progenitor cellen in de vervanging van insulinesecreterende B-cellen in preklinische modellen van type 1 diabetes. Period 1/1/2007-12/31/2009; annual direct cost: €30,000
- KUL CoE (Pi Verfaillie) Period: 11/1/2005 10/31/2009. Annual Direct Cost: €500,000/year
- Odvsseus Fund (PI Verfaillie) Period: 12/21/2006 12/31/2010. Annual Direct Cost: €1,390,000/year
- FP6-STREP: STROKEMAP (PI Verfaillie). Multipotent Adult Progenitor Cells to treat Stroke; Period: 10/1/2006 9/30/2009. Total Cost: €2,400,000; Total cost Verfaillie €420,000
- FP6-STREP: CHRYSTAL, (partner 6: Verfaillie) Cryobanking of stem cells for human clinical application. Period: 1/1/2007 12/30/2009. Total cost Verfaillie: €321,000
- SBO BRAINSTIM (PI Verfaillie); non-invasive imaging of stem cells in the brain; Period 10/1/2007-9/31/2011; Total Cost: €2.650.000; Total cost Verfaillie €550.000
- SBO: IMAGINE (Partner Verfaillie). Generation of improved paramagnetic particles for stem cell labeling and application in tumor therapy. Period: 1/1/2009 - 12/310/2013. Total cost Verfaillie: €250,000

PENDING FUNDING:

- FP7: BELISTEM (Partner Verfaillie). Stem cells suitable for liver regeneration. From the bench to the bed side; Period: 1/1/2010 - 12/30/2014. Total cost partner Verfaillie: €1.100.000
- FP7: STEMPER (Partner Verfaillie). Stem cells suitable for therapy of peripheral vascular disease. Period: 1/1/2010 -12/30/2014. Total cost KULeuven: €1,300,000
- S80: HEPSTEM (PI Verfaillie). Generation of mature hepatocytes from human induced pluripotency stem cells Period: 1/1/2010 - 12/310/2014. Total cost €2,500,000
- NIH GRANT 10121990 (PI Ekker, Co-I Verfaillie). Genomic Analysis of Hematopoietic Stem Cell Niche Formation, Maintenance & Function. Period: 1/7/2009 – 30/6/2014. Yearly Budget Verfaillie: \$125,000
- Dutch Diabetes Fund (PI P Devos, Co-I Verfaillie). Human fetal and adult progenitor cells as a source for insulin producing cells. 4/1/2009 31/3/2012. Total cost Verfaillie €250,000

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 Blood 79: 2821-2826. 1992. (Rapid communication) (IF: 10.9)
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Sequential Exposure to Cytokines Reflecting Embryogenesis: The Key for in vitro Differentiation of Adult Bone Marrow Stem Cells into Functional Hepatocyte-like Cells

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Differentiation of adult bone marrow stem cells (BMSC) into hepatocyte-like cells is commonly performed by continuous exposure to a cytokines-cocktail. Here, it is shown that the differentiation efficacy in vitro can be considerably enhanced by sequential addition of liver-specific factors (fibroblast growth factor-4, hepatocyte growth factor, insulin-transferrin-sodium selenite, and dexamethasone) in a time-dependent order that closely resembles the secretion pattern during in vivo liver embryogenesis. Quantitative RT-PCR analysis and immunocytochemistry showed that, upon sequential exposure to liver-specific factors, different stages of hepatocyte differentiation, as seen during liver embryogenesis, can be mimicked. Indeed, expression of the early hepatocyte markers alpha-fetoprotein and hepatocyte nuclear factor (HNF)3β decreased as differentiation progressed, whereas levels of the late liver-specific markers albumin (ALB), cytokeratin (CK)18, and HNF1\alpha were gradually upregulated. In contrast, cocktail treatment did not significantly alter the expression pattern of the hepatic markers. Moreover, sequentially exposed cells featured highly differentiated hepatic functions, including ALB secretion, glycogen storage, urea production, and inducible cytochrome P450-dependent activity, far more efficiently compared to the cocktail condition. In conclusion, sequential induction of the differentiation process, analogous to in vivo liver development, is crucial for in vitro differentiation of adult rat BMSC into functional hepatocyte-like cells. This model may not only be applicable for in vitro studies of endoderm differentiation but it also provides a "virtually unlimited" source of functional hepatocytes, suitable for preclinical pharmacological research and testing, and cell and organ development.

Key Words: bone marrow stem cells; hepatocytes; sequential differentiation; liver-specific growth factors; liver embryonic development; in vitro.

Drug development is aimed at identifying pharmacologically active drug candidates with a favorable toxicologic profile. The increasing number of safety criteria, imposed on newly designed molecules, leads nowadays to the urgent need of in vitro techniques in the industry, developed according to the principle of Russell and Burch. To date, several hepatocytebased in vitro models are available, however, they are not yet accepted into regulations, as they still require better characterization and optimization to reach the validation stage. Most primary hepatocyte cultures are in fact hampered by progressive occurrence of differentiation (De Smet et al., 2001; LeCluyse et al., 1996; Rogiers and Vercruysse, 1993). An alternative approach would be the use of postnatal progenitor/stem cells.

INTRODUCTION

Indeed, until recently, it was believed that tissue-specific stem cells could only differentiate into cells of the tissue of origin. However, a number of recent studies have suggested that adult stem cells may overcome germ lineage restrictions and express molecular characteristics of cells of different tissue origin, which has been termed "plasticity" (Jackson et al., 2001; Krause et al., 2001; Theise et al., 2000; Vourc'h et al., 2004). For example, hematopoietic cells may acquire characteristics of cardiomyocytes, cells of lung, gut, liver, blood vessels, skin, etc. (Jackson et al., 2001; Krause et al., 2001; Theise et al., 2000). This apparent plasticity can at least in some instances be explained by cell fusion (Wang et al., 2003). Other studies have described nonhematopoietic stem cells from bone marrow that are capable of differentiating in vitro in cells with mesodermal, ectodermal, and endodermal features (Jiang et al., 2002; Reves et al., 2001; Yoon et al., 2005). The mechanism through which these cells gain multipotency is not totally understood (Verfaillie, 2000). Multipotent adult progenitor cells, for instance, can be induced to express phenotypic and functional characteristics of hepatocytes; however, the degree of differentiation obtained till now is incomplete (Schwartz et al., 2002).

Therefore, in order to develop an in vitro model suitable for pharmaco-toxicological purposes, attempts were made here to

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optimize the differentiation efficiency of nonhematopoietic stem cells from bone marrow into functional hepatocytes.

Liver development is accomplished by a sequential array of biological events. Each step of cell growth and differentiation is tightly regulated by cell autonomous mechanisms and extracellular signals, including cytokines and growth factors. More specifically, during the initial phase of murine liver ontogeny (embryonic days [E] 8-9), fibroblast growth factors (FGFs), derived from adjacent cardiac mesoderm, commend the foregut endoderm to form the liver primordium (Duncan, 2000; Jung et al., 1999). During and after the mid-stage of hepatogenesis, surrounding mesenchymal cells secrete hepatocyte growth factor (HGF) and support as such the fetal hepatocytes (Kinoshita and Miyajima, 2002; Zaret, 2002). Around E11, the fetal liver becomes the major site for hematopoiesis. During this stage, hematopoietic stem cells produce oncostatin M that, in the presence of glucocorticoids, not only promotes fetal hepatic cell differentiation and maturation but also suppresses embryonic hematopoiesis. In contrast, oncostatin M alone fails to induce differentiated liver phenotypes, implying that glucocorticoids are essential triggers for hepatic maturation (Kinoshita and Miyajima, 2002; Schmidt et al., 1995; Zaret, 2002). In rodents, the final step of hepatic differentiation takes place several days after birth. The lack of terminal differentiation of primary hepatocytes in culture evidences that additional signals, probably generated through the extracellular matrix, are necessary (Kinoshita and Miyajima, 2002).

Here, the liver development was taken as exemplar to establish a culture model that more readily supports robust differentiation of bone marrow stem cells (BMSC) to mature hepatocyte-like cells. We compared two experimental setups: (1) BMSC were treated with a cockatial of liver-specific factors (FGF-4, HGF, insulin-transferrin-sodium selenite [ITS], and dexamethasone [Dex]) as previously described (Schwartz et al., 2002) or (2) innovative in this field, BMSC were exposed to a sequence of these compounds in a manner that closely reflects their temporal expression during in vivo hepatogenesis (FGF-4, followed by HGF, followed by a combination of HGF, ITS, and Dex).

MATERIALS AND METHODS

Isolation and culture of undifferentiates rat BMSC. BMSC were isolated from male Fisher rus (4-6 weeks old) and cultured as described by lings et al. (2002). Labava use def or expansion of BMSC included Corning 15 and 150 cm² tissue culture flasks, polystyene (both from VWR, Leuven, Belgium). Cell ksopyoping, neuroccioremal, and endotheil differentiation were determined as previously described (ling et al., 2002, 2003; Reys et al., 2000). Rath had cesses to food and water ad Bibtum and were bussed according to guidelines from the Institutional Animal Care and Use Committee of the University of Minnescota.

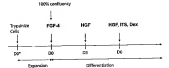
Hepatocyte differentiation. Rat BMSC from 60 population doublings on were used for differentiation into hepatocyte-like cells. BMSC were plated at 21 × 10³ cells/cm² on 1 mg/ml collagen type-1-coate culture plates and dishes (BD Falcon 24-well plate, polystyrene; BD Falcon 35 × 10 mm petri dishes,

polystyrene; NUNC F96 microwell plate, black, polystyrene; NUNC F96 microwell plate, clear, polystyrene [all from YWR]) in low-serum expansion medium (Glaug et al., 2002; Reyes et al., 2001). Once cells reached 100% confluence, they were wasted with basal medium (Biang et al., 2003) supplemented with 0.03mM includimation, 0.25mM sodium-privates and 1.623mM glutamine (all from Sigma, Bornera, Beglium). Subscepensly, cells were caltured in the presence of liver-specific cytokines and growth factors, added either as a cockrail (basal medium + 10 ng/m) FGF-4, 20 ng/ml fBGF [all from R8D Systems, Minnespola, MN]. X ITS and 20 up IDs Ex [all from Sigma]) or sequentially (days) 0-3: basal medium + 10 ng/ml FGF-4 (days) 3-6: basal medium + 20 ng/ml FGF-1 (14) FS (

Quantitative RFPCR: For PCR analysis, 1 ug RNA was reverse transcribed to DNA using Superactify III evener transcriptes and random hazamer primers (navingean, Marcibeke, Belgium). The resulting RF products were essentially supplied as proviously described (lang et al., 2003; Schwart et al., 2002). Three extra steps were included to ensure the purity of the PCR products: 99°C for 15's, 60°C for 20's, and 99°C for 15's. The primers used for amplification and the products expected are described in (Jining et al., 2003; Schwart et al., 2002). The RNA levels were normalized using 18S and compared with the RNA levels in undifferentiated BMSC (negative control) and freshly isolated primary art hespotsets (positive control). As a negative control for the primers; a no template oDNA-PCR reaction was run under the same conditions. The authenticity and size of the PCR products were contimor by melting curve analysis (using software provided by Perich Eliner, Lennik, Belgium) and gel electrophoresis.

Immunocytochemistry. Differentiated BMSC were fixed either with thand for 10 min at = 20°C (cytoskeletal proteins) or with 4% part-formalichyted (Electron Microscopy Sciences, Fort Washington, PA) for 10 min at 4°C, followed by incubation with 10 mM glycin to saturate reactive groups (nuclear and cytophamic markers). The fixed cells were permeabilized for

A) Sequential exposure to liver-specific factors



B) Exposure to a cocktall of liver-specific factors

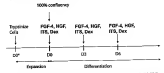


FIG. 1. Schematic presentation of the differentiation protocol. BMSC, at 100% confluency, were exposed either sequentially (A) or simultaneously (B) to liver-specific factors, D0°, day that BMSC were plated at 21 × 10° cells/cm° on collagen type I in low-serum expansion: medium.

15 min with 0.1% Triton in phosphate-buffered saline (Electron Microscopy Sciences) and blocked for 30 min with 1% bovine serum albumin/5% donkey serum block buffer at room temperature. After blocking, cells were incubated overnight at 4°C with primary antibody (fluorochrome-conjugated or nonconjugated) and washed three times with phosphate-buffered saline. In case the primary antibody was not conjugated, cells were incubated for 2 h at room temperature with secondary fluorochrome-conjugated antibody. After incubation, slides were washed again with 0.1% Triton in phosphate-buffered saline and mounted with Vectashield with DAPI (Vector Laboratories, Burlingame, CA). As a negative control, cells were incubated with appropriate gamma immunoglobulines (Jackson Immunoresearch, Cambridgeshire, UK) and immunostained under the same conditions. In order to evaluate the localization of cytochrome P450 (CYP) proteins, mitochondria and endoplasmic reticulum were counterstained with the carbocyanine dye DiOC6 (Molecular probes, Invitrogen). Cells were analyzed using fluorescence microscopy with a Zeiss Axiovert scope. To enumerate the number of cells expressing a given marker, all nuclei of positive-stained cells were counted and compared to the total number of cells evaluated. The primary antibodies against alpha-fetoprotein (AFP) (goat), hepatocyte nuclear factor (HNF)3β (goat), and HNFIα (rabbit) were purchased from Santa Cruz, (Heidelberg, Germany). Anti-cytokeratin (CK)18 (mouse, FTTC-conjugated) and anti-albumin (ALB) (goat, FTTC-conjugated) antibodies were from Sigma and Bethyl Laboratories (Montgomery, TX), respectively. The antibodies against CYP1A1 and CYP2B1/2 (both goat) came from Dajichi pure chemicals, BD Biosciences (Tokyo, Japan). Respective secondary antibodies were purchased from Jackson Immunoresearch.

Albumin ELISA. ALB concentrations, secreted into the culture media, were analyzed by ELISA (Koebe et al., 1994).

Urea usags. The produced urea concentrations were, after 24-be exposure of the cells to Gind NHLCI, colonomically measured in culture media according to the manufacturer's instructions Quantichrom Urea sassy list. Biosassy Systems, Brussels, Helgium). Petch culture media supplemented with 6mM NHLCI and 4b-cultured adult rat hepatocytes were used as a negative and ossitive control. It is executively.

Glycogen storage. Intracellular glycogen was analyzed by Periodic-acid-Schiff staining (PAS-kit 395B-1KT, Sigma) according to the manufacturer's

instructions. Amyloglucosidase (Sigma)-treated cells and 4 h-cultured adult rat hepatocytes were used as a negative and positive control, respectively.

Alkozyresorufin-O-dealkylase assay. Ethoxyresorufin-O-deethylase (EROD) and pentoxyresorufin-O-dealkylase (PROD) activities were assessed as previously described (Denato et al., 1993) with some minor modifications: in our setup, cells were incubated with 20µM 7-ethoxyresorufin and 18µM 7-entoxyresorufin (all from Signan) for 30 min.

To evaluate the industibility of CVPDB1/2 and CVP1A1/2, respectively, cells were, after 24 days of differentiation, exposed to phenobarthial (BB, final concentration 1mM) and 3-methylcholanterne (MC, final concentration 2µM; all from Sigma). Media, supplemented with either PB or MC, were diversely renewed from that time on. Fresh culture media and 4 h-cultured adult rat henotocytes were used as a negative and positive control, respectively.

Statistics. Results are expressed as mean \pm SD. Statistical analyses were performed using one-way ANOVA and Student's t-test. The significance level was set at 0.05.

RESULTS

Characterization of the Differentiation Pattern of Rat BMSC into Hepatocyte-like Cells: Sequential versus Cocktail Exposure

Morphological Features

Previously, it has been shown that BMSC could differentiate into hepatocyte-like cells upon simultaneous exposure to a mixture of well-defined cytokines and growth factors (Schwartz et al., 2002). However, using this approach, a rather heterogeneous population of epithelioid cells and other cell types was obtained. Moreover, no polygonal-shaped cells and only few binucleated cells were formed (Fig. 2). In an attempt

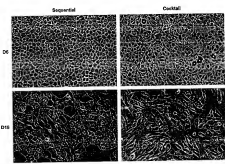


FIG. 2. Light-microscopic analysis of BMSC-derived hepatocyte-like cells upon sequential or simultaneous exposure to liver-specific factors at days 6 and 18. Original magnification of 20 × 10, phase contrast.

to improve the differentiation of nonhaematopoietic stem cells from bone marrow into hepatocyte-like cells, BMSC were exposed to the same well-defined hepatogenic factors, but in a sequential way. More specifically, cytokines and growth factors were added at defined points in time, in a manner that closely resembles the in vivo process of embryonic liver development as specified in "Materials and Methods" section (Duncan, 2000; Jung et al., 1999; Kinoshita and Miyajima, 2002; Schmidt et al., 1995; Zaret, 2002), in this novel setup, epithelioid cells appeared in culture from day 6 on (Fig. 2). However, at that moment these cells were still surrounded by spindle-shaped cells. After 14 days, less fibroblastic cells were seen and some binucleated cells appeared. After 18 days, most cells exhibited a polygonal shape (Fig. 2).

Characterization at the Molecular Level

In a next set of experiments, we evaluated whether these morphological differences were associated with distinct pattens of differentiation at the molecular level. Therefore, the expressions of early (AFP and HNF3B) and late (ALB, CK18, and HNF1a) liver-specific markers were analyzed at both the mRNA (Fig. 3) and protein levels (Figs. 4 and 5).

mRNA expression. In both sequential and cocktail culture conditions, AFP, HNF3β, ALB, CK18, and HNF1α were

expressed in a time-dependent manner during BMSC differentiation. Both the pattern and the level of expression, however, differed considerably between the culture methods. In fact, upon sequential exposure to liver-specific factors, maximal AFP mRNA expression occurred after 6 days (Fig. 3), 4 days later than seen in the cocktail condition, but was 1.2-fold higher than the maximal level observed in cocktail-exposed cells. AFP mRNA expression disappeared completely in both conditions by day 11 of culture. In sequentially treated cells, downregulation of AFP mRNA expression was nicely followed by a second transient, though more pronounced, induction of the early liver-specific marker HNF3 \beta as well as by a steady upregulation of the late hepatic markers ALB, CK18, and HNF1α (Fig. 3). More specifically, HNF3β mRNA expression started at day 2, reached maximal levels at day 10 and decreased rapidly thereafter (Fig. 3). ALB and CK18 mRNA expression, on the other hand, gradually increased from days 4 and 10 of culture, respectively, until maximal levels were reached at day 18 (Fig. 3). In sharp contrast to these observations, changes in HNF3B, ALB, and CK18 mRNA levels were negligible upon simultaneous exposure to all hepatogenic factors (p < 0.001; one-way ANOVA). In addition, the mRNA of the late liver-specific marker ALB remained very low in cocktail-exposed BMSC, suggesting an immature hepatic differentiation status. Finally, upon sequential exposure, HNF1a

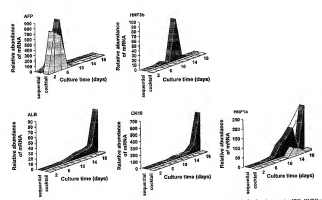


FIG. 3. Analysis of hepsuccyte differentiation at the mRNA level (abundance in cultured cells relative to freshly isolated as hepsuccytes [85]). BMSC were either sequentially or simultaneously exposed to liver-specific factors. Values represent means of three independent experiments. The dotted lines represent monested time points. They are assumptions, based on the existing expression pattern.

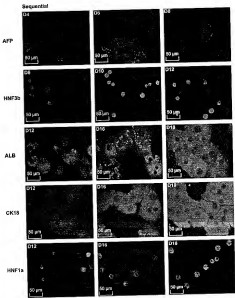


FIG. 4. Characterization at the protein level of BMSC differentiation into hepatocyte-tike cells upon sequential exposure to liver-specific factor. Immunocytechemistry was performed for AFP-03, IMF3β-03, CM3-FITC, IMF3 e-03, and ALB-FITC. Nuclear counterstanking was actionable using DAFI. Original magnification of ×230. Seel but, 30 μm. Suitings shown have the same magnification and are representable for at least five separate experiments.

mRNA expression gradually increased from day 6 on whereas in the cocktail condition, HNF1a mRNA induction was delayed by 4 days and occurred only transiently (Fig. 3). Moreover, maximal levels, obtained at day 12, were about twofold lower than the levels observed in 12-day-old BMSC in the sequential condition. Thus, sequentially exposed BMSC underwent a consecutive array of developmental stages comparable with in vivo hepatogenesis while exposure to a cockail of cytokines and growth factors induced an aberrant expression pattern of differentiation when compared to liver embryogenesis.

Protein expression. In order to support the results obtained at the mRNA level, immunocytochemistry analyses were performed in parallel (Figs. 4 and 5). After 4 days of differentiation, cells expressed AFP, regardless of the experimental setup (Figs. 4 and 5). After expression occurred only transiently in both conditions (Figs. 4 and 5) and was undetectable by day 12 of culture (data not shown). Upon sequential exposure to liver-specific factors, a maximal positive staining of HNF3β (92 ± 8%) was noticed at day 10, leveling off thereafter (Fig. 4). Treatment with all factors

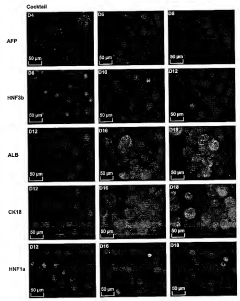


FIG. 5. Characterization at the protein level of BMSC differentiation into hapatocyte-like cells upon simultaneous capouse to liver-specific factors. Immunocytechemistry was performed for AFP+03, INIV3B+03, CMS+HTC, INIV1s+03, and ALB-HTC. Nuclear counterstaining was assessed using DAFI, Original magnification of X230, Sea they, 50 pm. Statistical shows have the same magnification and are representative for at least five long-rate experiments.

simultaneously, however, revealed no more than $24 \pm 7\%$ INNF3β-positive cells throughout the culture period (Fig. 5). As differentiation progressed, extensively increased stainings for ALB, CK18, and HNF1a were detected upon sequential exposure to cytokines and growth factors, in accordance with the results obtained at the RNA level (Figs. 3 and 4). Consequently, after 18 days, $92 \pm 2\%$, $94 \pm 3\%$, and $89 \pm 9\%$ of the cells, respectively, stained positive for these markers (Fig. 4), which is in sharp contrast to only $32 \pm 4\%$, $63 \pm 5\%$, and $22 \pm 4\%$ of the cocktail-exposed cells, respectively (p < 0.001; Student's f-test) (Fig. 4).

In addition, in order to state the immunocytochemistry data with certainty, immunoblotting has been performed in parallel once (data not shown). In line with the previous results obtained at both the mRNA and protein level, sequentially exposed cells expressed liver-specific proteins more abundantly than cells in the cocktail setup. However, since this approach consumes large numbers of cells, i.e., at least 25–50 µg of protein is needed to analyze one liver-specific marker at one point in time, the analysis was not repeated. Alternatively, as measuring CYP activity (the set of EROD/PROD) and their inducibility are widely accepted as final end point to evaluate the suitability of

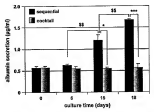


FIG. 6. ALB secretion in sequentially—and cocktail-exposed BMSC. The results shown are representative for five independent experiments, seek performed in duplicate.** ****. ALB-secretion significantly differs among sequentially—and cocktail-exposed BMSC with ρ < 0.003 and ρ < 0.001, respectively (Student's *text). SS: ALB-secretion in sequentially exposed BMSC is significantly upgesquated from day 15 on with ρ < 0.01 (Student's *text).

cells as in vitro models for pharmaco-toxicological screening of drugs (De Smet et al., 2001; Donato et al., 1993, 2001; LeCluyse et al., 1996; Rogiers and Vercruysse, 1993), we enlarged, in a next set of experiments, the data set on cell functionality in order to increase confidence in our data.

Hepatic Functionality

In order to assess whether these hepatocyte-like cells derived from the bone marrow also acquired typical functional hepatic features, ALB secretion, ammonia metabolism, glycogen storage, expression of CYP proteins in parallel with their activity and inducibility were evaluated.

ALB secretion. Sequentially treated BMSC significantly upregulated the ALB secretion rate from day 15 onward (p < 0.01, Student's t-test) (Fig. 6). On the contrary, BMSC exposed to a cocktail of liver-specific factors did not secrete ALB above basal levels, corresponding to 0.55 µg/ml (Fig. 6).

Ureogenesis. Upon sequential exposure to hepatogenic factors, the urea production increased over culture time, reaching adult levels after 30-33 days. In contrast, cocktail-exposed cells synthesized, even at peak production, 24% significant lower urea levels (p. 0.05; Student's 1-test) (Fig. 7).

Glycogen storage. Furthermore, upon sequential treatment with cytokines and growth factors, glycogen uptake was first seen after 21 days of culture, 6 days earlier than in the cocktail condition. After 30 days of culture, about 86% of the cells stored glycogen, regardless of the culture method (Fig. 8).

CYP protein-expression, activity, and inducibility. In the sequential setup, phase I CYP1A1 and CYP2B1/2 proteins were expressed within and nearby the endoplasmic reticulum and mitochondria (Fig. 9). The level of expression gradually

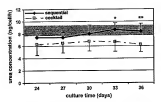


FIG. 7. Urea production in sequentially- and cockial-exposed BMSC. Gray area represents urea levels, produced by h -builtured adult hepatocytes. The graph is representative for four separate experiments, each performed in duplicate. \star *** Use production significantly differs among sequentially- and cockial-exposed BMSC with p < 0.05 and p < 0.01, respectively (Student's 4-text).

increased as differentiation progressed. After 30 days, 78 ± 1 and $79 \pm 3\%$ of the cells stained positive for CYP1A1 and CYP2B1/2, respectively (Fig. 10). In sharp contrast to these observations, cocktail-exposed cells only showed modest CYP expression over culture time (Figs. 9 and 10).

In addition, we investigated whether CYP1A1 and 2B1/2 were functionally active by measuring the respective EROD and PROD activities in both conditions (Figs. 11 and 12).

In line with the results found at the protein level, sequentially exposed cells exhibited markedly higher EROD and PROD activity rates compared to the cocktail model (p < 0.05 at days 36 and 39; Student's r-test) (Figs. 11 and 12). Upon sequential exposure to live-specific factors, a transient foundfol finerase in PROD activity was displayed by days 27–30, approaching the level of 4h-cultured adult rat hepatocytes, versus only a twofold increase after cocktail treatment (Fig. 11). In addition, in the former setup, EROD activity gradually increased from days 27 to 36 towards levels measured in 4h-cultured adult rat hepatocytes, whereas CYP1A1/2-dependent activities appeared only transiently in cocktail-exposed cells between days 30 and 33 and declined to lambs to modetectable levels on days 36 (Fig. 12).

CVP-inducibility is considered as the most representative metabolic function of the adult hepatic phenotype (Gomez-Lechon et al., 2004; Rogiers and Vercruysse, 1993). Therefore, the responsiveness of both CYP1A1/2 and CYP2B1/2 to their respective prototype inducers MC and PB was analyzed in parallel. PROD activities were induced up to 1.4-fold after 6-day exposure to PB (i.e., on day 30), regardless of the experimental setup (Fig. 11). The inducibility persisted for 6 days in sequentially exposed cells but not in the cocktail condition. A significant CYP1A1/2-dependent response to MC was observed on days 36-39 (p < 0.001 and p < 0.01 at days 36 and 39, respectively; Student's *Lest*) in the sequential model. Conversely, MC barely induced EROD activities upon culture with all liver-specific factors simultaneously (Fig. 12).

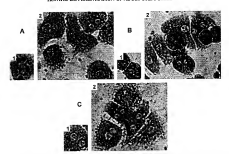


FIG. 8. Glycogen storage in 30-day-old sequentially-(A) and cocktsil-exposed BMSC (B) and 4 h-cultured adult rat begatocytes (C), in the presence (I) and absence (2) of anyloglucosidase, respectively, Glycogen and nuclei are colored magenta and blue, respectively. Original magnification of ×400. Stainings shown are representative for three separate coperiments.

DISCUSSION

In recent years, adult-derived stem cells have become a hot topic in the field of molecular, cellular, and clinical biology, as well as in pharmaco-toxicology. Indeed, stem cells have an extensive self-renewing potential and many of them are considered multipotent (Jackson et al., 2001; Krause et al.,

2001; Theise et al., 2000; Vourc'h et al., 2004). This interest in adult stem cells has in particular been triggered by the numerous ethical dilemmas surrounding the use of embryonis stem cells in preclinical and clinical research (Henningson et al., 2003; McLaren, 2001). The best-characterized stem cell compartment is the bone marrow consisting of two stem cell populations, referred to as the hematopoietic and the messenchymal

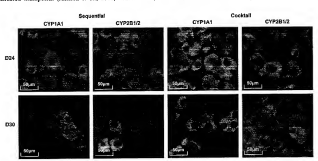


FIG.9. Detailed view of expression of phase Islociansformation enzymes at days 24 and 30 upon sequential or simultaneous expanse to Piver-specific Rotors. Immunoscytochemistry was performed for CYPIA1-e-93 and CYPE316/2-93. Mitochondria and endoplasatic reinclum were counterstand with a green fluorescent carbocyraine. Nuclei were counterstanded with DAPI. Original magnification of x320. Scale bar, 30 µm. Stainings shown have the same magnification and are representative for three separate experiments.

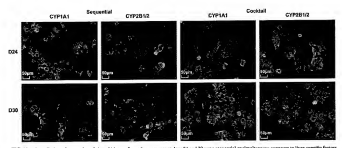


FIG. 10. Overall view of expression of phase I biotransformation enzymes at days 24 and 30 upon sequential or simultaneous exposure to liver-specific factors. Immunocytochemistry was performed for CYPIAI-cy3 and CYPIBIA2-cy3. Mitochondria and endoplasmic reticulum were counterstained with a green fluorescent eurobcysanie. Nuclei were counterstained with DAPI. Magnification of ×112. Scale bar, 50 µm. Stainings shown have the same magnification and are representative for three separate experiments.

stem cells (Huttmann et al., 2003). Previously, Schwartz et al. (2002) described a population of cells in postnatal rat bone marrow, copurified with mesenchymal stem cells, that were capable of differentiating into cells of endodermal (hepatocytes) origin upon exposure to well-defined hepatogenic factors. These culture conditions yielded, however, a mixture of epithelioid

cells and other cell types. Therefore, attempts were made here to improve the hepatic differentiation process through exposure of BMSC to the same liver-specific factors in a sequential time-dependent manner, reflecting their secretion during in vivo hepatogenesis (Duncan, 2000, Jung et al., 1999; Kinoshita and Miyajima, 2002; Schmidt et al., 1995; Zaret, 2002).

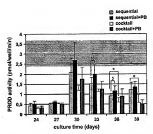


FIG. 11. PROD activities and responsiveness to InM TB in differentiated BMSC upon sequential or simultaneous exposure to liverspecific factors. FB was added dually, starting on day 24. Gray area represents PROD activity measured in untreated 4 h-cultured abilit rat hepatocytes. The graph is representative for two separate experiments, each, performed in diplicate. *: PROD activity significantly differs among sequentially- and cocktuil-exposed BMSC with p < 0.05 (Squards + leaf), 3.5; PB significantly induced PROD activity of sequentially-exposed BMSC with p < 0.05 and p < 0.01, respectively (Sudent's + least).

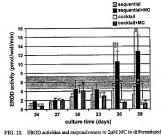


FIG. 2. EMD/8 dytes and resemble to the representation to Just nc. in Universities MSC upon sequential or simulations exposure to Personage Continue Move was added fully, marked a challengt and the properties of the properties. The properties of the properties

Under these culture conditions, BMSC acquired morphological features (polygonal-shaped and binucleated cells) similar to those of primary hepatocytes (Ferrini et al., 1997; Katsura et al., 2002). Furthermore, more than 85% of these epithelioid cells expressed liver-associated genes and proteins (AFP, HNF3B, ALB, CK18, and HNF1a) in a comparable timedependent manner as observed during in vivo liver embryogenesis, Indeed, AFP expression is first detected in embryonic endoderm around E8.5 (Cascio and Zaret, 1991) and precedes ALB and HNF1\alpha expression, detected around E9.5 and E10.5, respectively (Ott et al., 1991; Shiojiri, 1981). This finding implicates that, in this setup, the BMSC differentiation process could serve as a model of early mammalian endoderm differentiation. In contrast, upon exposure to a cytokine/growth factors-cocktail, the expression patterns differed from the normal sequence seen during in vivo hepatogenesis as HNF1 a expression preceded that of ALB. Indeed, HNF1 a is only expressed in fully differentiated cells and not in un- or dedifferentiated cells (Cereghini et al., 1988), as was noticed here upon cocktail treatment. In addition, significantly lower levels of liver-specific markers were expressed. The higher levels of ALB and CK18 expression in the sequential condition are probably due to the higher levels of both the early (HNF3ß) and late (HNF1a) transcription factors. It is well documented that liver-enriched transcription factors act cooperatively and synergistically to promote liver-specific gene transcription (Cereghini et al., 1992; Darlington, 1999; Duncan, 2000; Hayashi et al., 1999; Shim et al., 1988). In this regard, it was previously shown that HNF3 B positively regulates the expression of HNF4α and HNF1α (Darlington, 1999; Duncan et al., 1998). Furthermore, it is believed that HNF3β serves as the initiator of a cascade of regulatory events resulting in endoderm induction (Ang et al., 1993; Darlington, 1999; Duncan, 2000; Levinson-Dushnik and Benvenisty, 1997). Hence, the minor changes in HNF3B expression levels in the cocktail condition may only result in low levels of ALB and CK18 transcripts and protein.

The initiation and induction of AFP expression is not yet completely understood. It can be assumed that additional factors are involved in its transcriptional activation, as in both culture conditions, only minimal levels of HNF3 β were detected at the time of AFP expression. Further research will be needed to fully elucidate the transcriptional hierarchy mediating differentiation of BMSC toward hepatocytes.

The presence of both morphologic and phenotypic features, similar to that of primary hepatocytes, does, however, not fully prove the differentiation of BMSC into mature hepatocytes. Indeed, during the terminal step of liver organogenesis, the liver becomes a functional and metabolic organ, performing an essential role as detoxifying center of the body (Gomez-Lechon et al., 2004; Kinoshita and Miyajima, 2002; Zaret, 2002). Interestingly, functional maturation occurred in both experimental setups, but to a different extent. Hepatic metabolic functions, including ALB secretion, urea production,

storage of glycogen, and CYP-activity/inducibility, were manifested most prominently upon sequential exposure to hepatogenic factors. Under these culture conditions, ALB secretion was in fact significantly upregulated to levels comparable to those obtained in both 2- to 7-day-old immobilization and 7-day-old monolayer cultures of primary rat hepatocytes. The latter measurements are performed on a regular basis in our laboratory (Beken et al., 2001; Vanhaecke et al., 2004). In addition, both the urea production and EROD/PROD activities reached levels comparable to 4 h-cultured primary rat hepatocytes. Response to prototype inducers was as expected: pronounced upon exposure to MC and discrete upon PB treatment. The level of induction, however, remained lower in comparison to cultured adult rat hepatocytes. More specifically. EROD activity increased up to fourfold after 15-day exposure to MC in sequentially exposed BMSC versus maximal sevenfold in 2-day treated rat hepatocytes (Donato et al., 1993). Nevertheless, to our best knowledge, this is the first time that EROD (CYP1A1/2) activity/inducibility is demonstrated in hepatocyte-like cells derived from BMSC.

The less mature phenotype of cocktail-exposed cells could possibly be ascribed to altered and lower expression of HNFtype liver-enriched transcription factors in this setup. Experiments using hepatoma cell lines and HNF-null mice have in fact demonstrated the important role of HNFs in the regulation of genes that are involved in biotransformation (Cyps) and ammonia metabolism (ornithine-transcarbamylase gene) (Gomez-Lechon et al., 2004; Inoue et al., 2002; Rodriguez-Antona et al., 2002). Similar to the results reported here, inducible CYP2B1/2-activity was also found by Schwartz et al. (2002) after exposure to a cocktail of the same cytokines and growth factors, although at an earlier time in culture. Some variation in time-specific gene and protein expression could probably be attributated to intraspecies differences and subtle changes in the differentiation procedure (i.e., type of culture plate coating, serum, etc.).

In summary, during the first 18 days of the hepatic differentiation process of BMSC, cells, and sequentially exposed BMSC in specific, underwent a sequential array of developmental stages, characterized by the down- and upregulation of early and late liver-specific markers, respectively. As differentiation progressed, i.e., from day 18 onward, expression of mature hepatic markers persisted at steady levels (data not shown) and cells gradually underwent functional hepatic maturation. In specific, sequentially treated BMSC accomplished hepatic functions at levels comparable to those of primary rat hepatocytes, cultured for 4 h to 2 days. Our results thus clearly show that a more pronounced and homogeneous differentiation of BMSC into functional hepatocyte-like cells can be obtained by sequentially directing the differentiation process analogous to liver embryogenesis. Moreover, differentiation appears to occur via steps commonly defined for in vivo endodermal lineage specification and subsequent hepatocyte differentiation and maturation. Further investigations, in order 340 SNYKERS ET AL.

to elucidate the molecular mechanisms underlying the changes described herein, are underway.

This model opens new perspectives: it may not only be applicable to study endoderm differentiation in vitro but it also offers the possibility to purify and culture multipotent stem cells from nonembryonic origin as an unlimited cell source for pharmaco-toxicological research and testing, and cell and organ development. It might even open a road to trigger cell fate and "trans" differentiate uncommitted cells from different tissues towards endodermal lineages.

SUPPLEMENTARY DATA

Supplementary data are available online at http://toxsci. oxfordiournals.org/.

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